# Neoplasia Lecture 3

**CARCINOGENESIS** 

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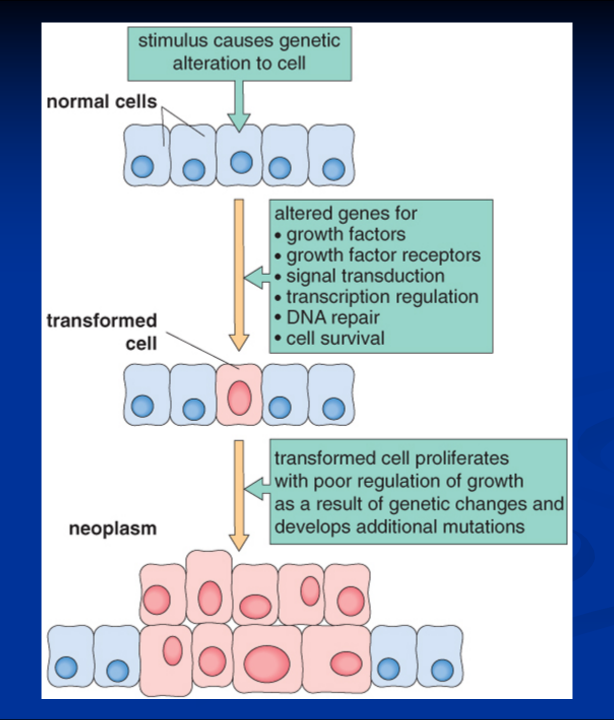
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Foundation block 2014 Pathology

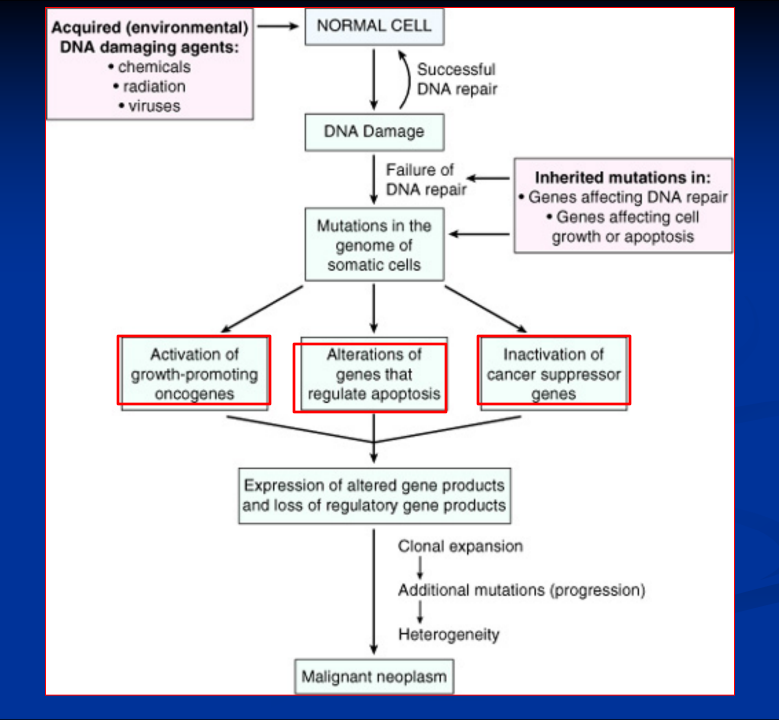
### **CARCINOGENESIS**

- Carcinogenesis is a multistep process at both the phenotypic and the genetic levels.
- It starts with a genetic damage:
  - Environmental
    - Chemical
    - Radiation
    - Infectious
  - Inhereted

- Genetic damage lead to "mutation"
- single cell which has the genetic damage undergoes neoplastic proliferation (clonal expansion) forming the tumor mass



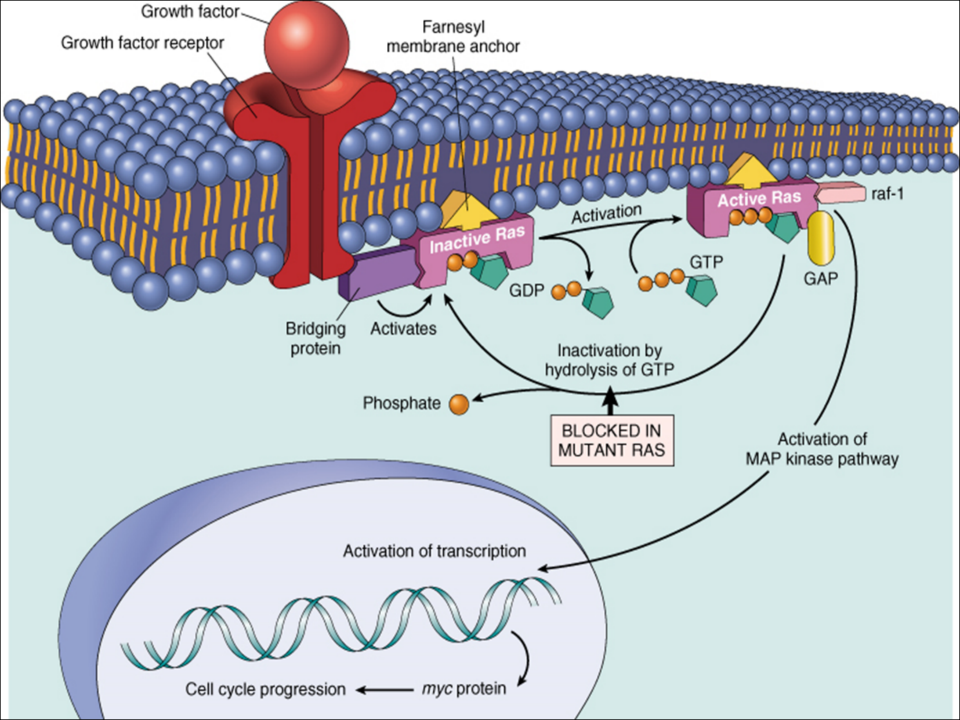
- Where are the targets of the genetic damage??
- Four regulatory genes are the main targets:
  - Growth promoting protooncogenes
    - Protooncogene > mutation > oncogene
  - Growth inhibiting (supressors) genes
  - Genes regulating apoptosis
  - DNA repair genes



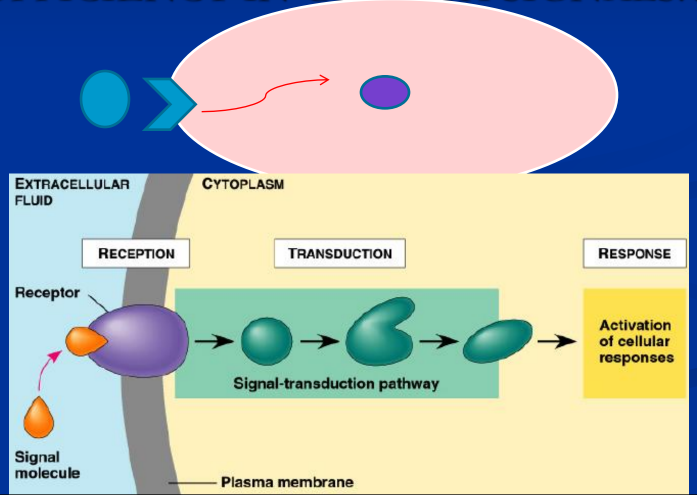
- Main changes in the cell physiology that lead to formation of the malignant phenotype:
  - Self-sufficiency in growth signals
  - Insensitivity to growth-inhibitory signals
  - Evasion of apoptosis
  - Limitless replicative potential
  - Sustained angiogenesis
  - Ability to invade and metastsize

- A Self-sufficiency in Growth signals:
  - Oncogene: Gene that promote autonomous cell growth in cancer cells
  - They are derived by mutations in protooncogenes
  - They are characterized by the ability to promote cell growth in the absence of normal growthpromoting signals
  - Oncoproteins : are the products

- Remember the cell cycle !!
  - Binding of a growth factor to its receptor on the cell membrane
  - Activation of the growth factor receptor leading to activation of signal-transducing proteins
  - Transmission of the signal to the nucleus
  - Induction of the DNA transcription
  - Entry in the cell cycle and cell division

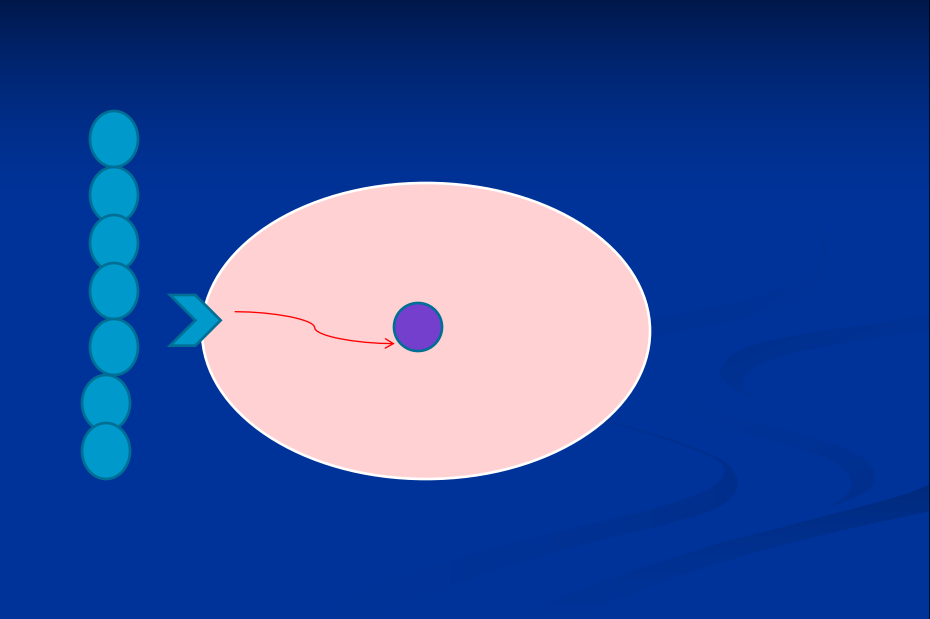


HOW CANCER CELLS ACQUIRE SELF-SUFFICIENCY IN GROWTH SIGNALS??



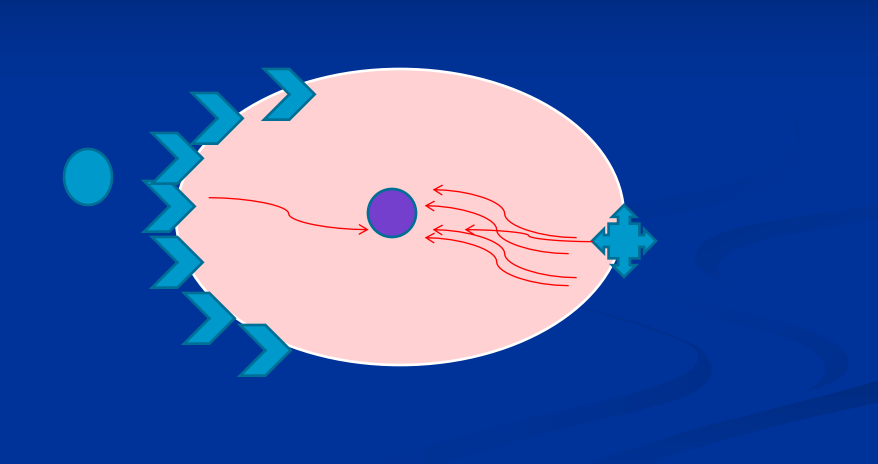
#### 1- Growth factors:

- Cancer cells are capable to synthesize the same growth factors to which they are responsive
  - E.g. Sarcomas ---- > TGF-α
    Glioblastoma----> PDGF



### 2-Growth factors receptors:

- Receptors --- mutation ---- continous signals to cells and uncontroled growth
- Receptors --- overexpression ---cells become very sensitive ----hyperresponsive to normal levels of growth factors

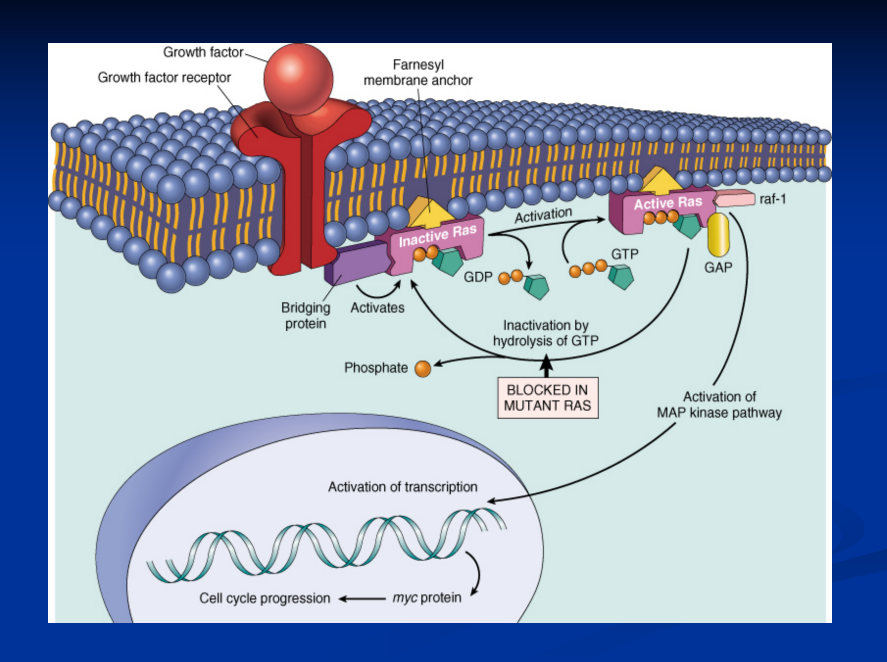


- Example:
  - Epidermal Growth Factor (EGF) Receptor family
    - HER2
      - Amplified in breast cancers and other tumors
      - High levels of HER2 in breast cancer indicate poor prognosis
      - Anti- HER2 antibodies are used in treatment

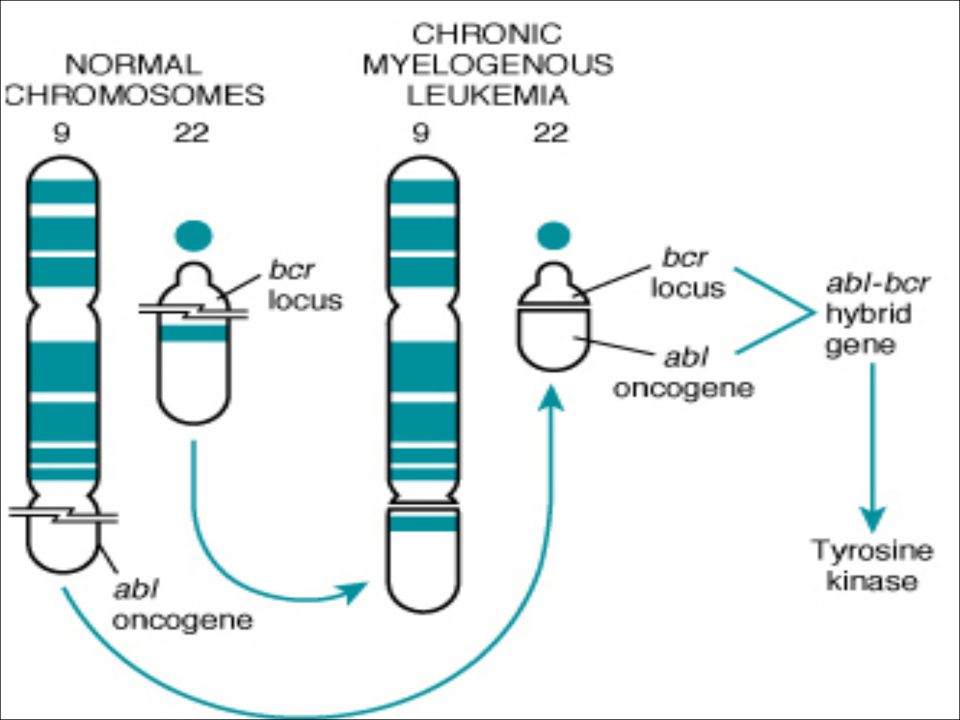
- 3- Signal-transducing proteins:
- They receive signals from activated growth factors receptors and transmitte them to the nucleus. Examples:
  - RAS
  - ABL

#### RAS:

- 30% of all human tumors contain mutated RAS gene . E.g : colon . Pancreas cancers
- Mutations of the RAS gene is the most common oncogene abnormality in human tumors
- Mutations in RAS --- cells continue to proliferate

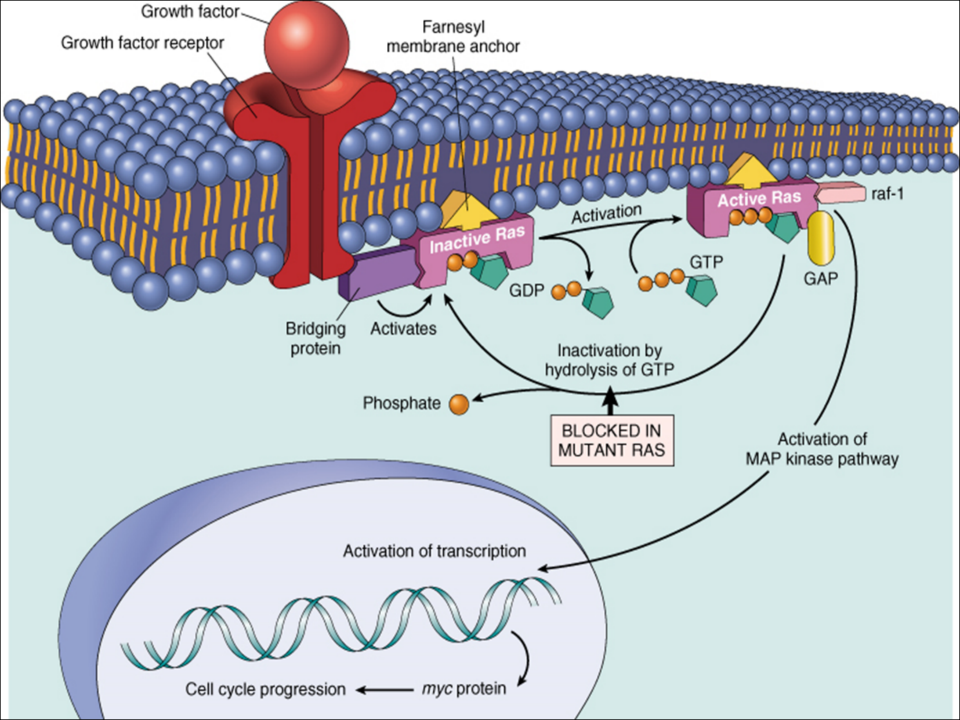


- ABL gene
  - ABL protooncogene has a tyrosine kinase activity
  - Its activity is controlled by negative regulatory mechanism
  - E.g.: chronic myeloid leukemia (CML):
    - t(9,22) ---ABL gene transferred from ch. 9 to ch. 22
    - Fusion with BCR ---> BCR-ABL
    - BCR-ABL has tyrosine kinase acttivity --- (oncogenec)



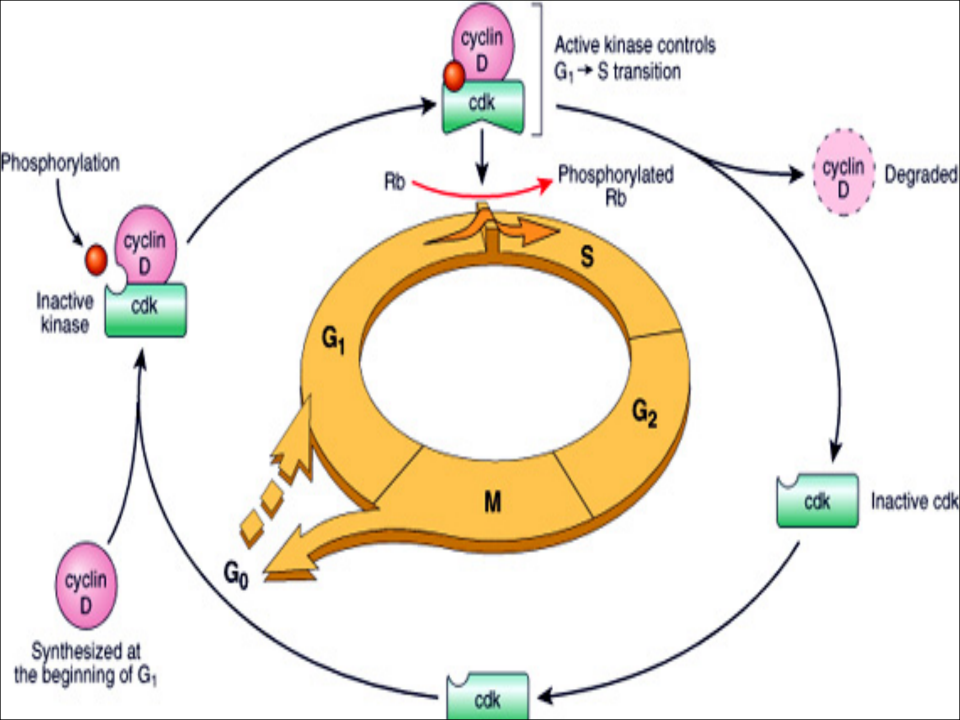
■ CML patients are treated with (Gleevec) which is inhibitor of ABL kinase

- 4- Nuclear transcription factors:
  - Mutations may affect genes that regulate transcription of DNA → growth autonomy
  - E.g. MYC
    - MYC protooncogene produce MYC protein when cell receives growth signals
    - MYC protein binds to DNA leading to activation of growth-related genes



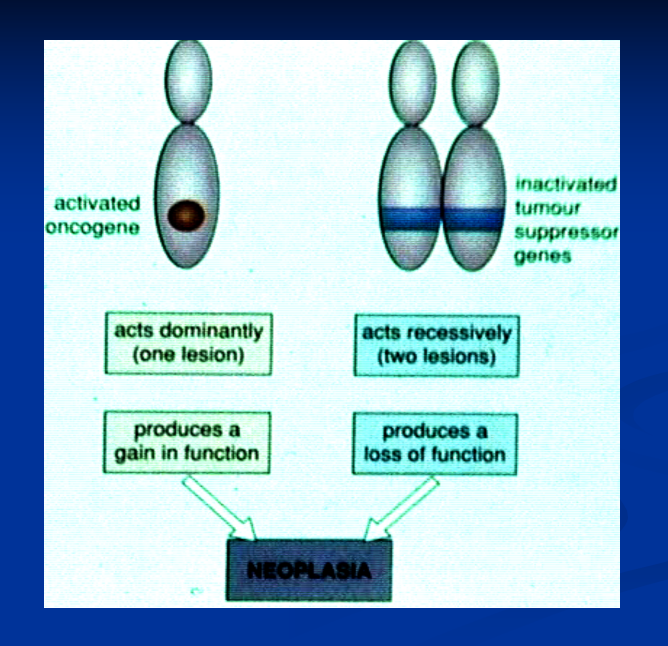
- Normally ... MYC decrease when cell cycle begins ...but ..in tumors there is sustained expression of MYC → continuous proliferation
- E.g. Burkitt Lymphoma; MYC is dysregulated due to t(8,14)

- 5- Cyclins and cyclins- dependent kinases (CDKs)
  - Progression of cells through cell cycles is regulated by CDKs after they are activated by binding with cyclins
  - Mutations that dysregulate cyclins and CDKs will lead to cell proliferation ...e.g.
    - Cyclin D genes are overexpressed in breast, esophagus and liver cancers.
    - CDK4 is amplified in melanoma and sarcomas



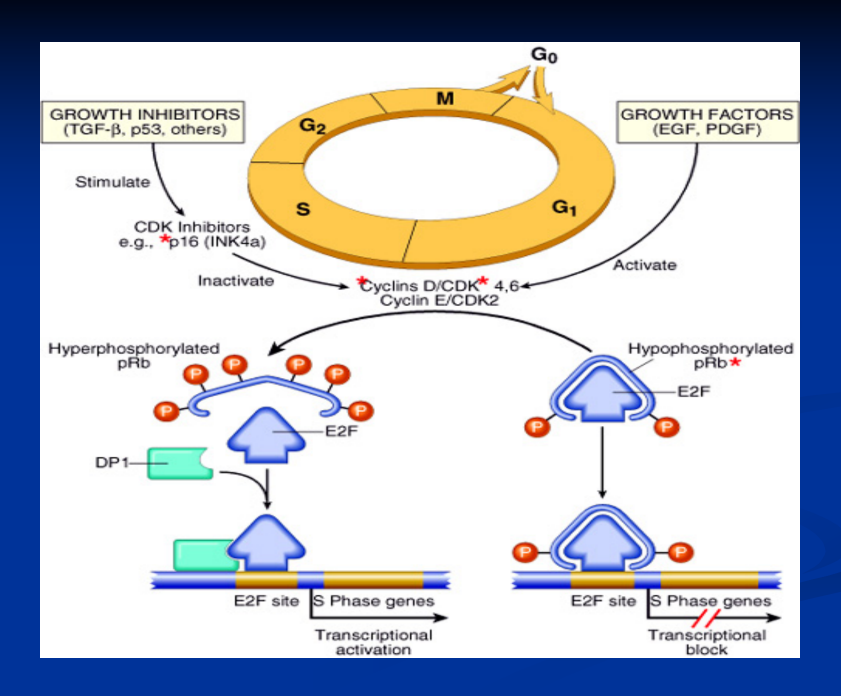
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  - C- Evasion of apoptosis
  - D- Limitless replicative potential
  - E- Sustained angiogenesis
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- 2. Insensitivity to growth-inhibitory signals
- Tumor suppressor genes control (apply brakes) cells proliferation
- If mutation caused disruption to them → cell becomes insensitive to growth inhibition → uncontrolled proliferation
- Examples: RB, TGF-β, APC, P53



- RB (retinoblastoma) gene:
  - First tumor supressor gene discovered
  - It was discovered initially in retinoblastomas
  - Found in other tumors, e.g. breast ca
  - RB gene is a DNA-binding protein
  - RB is located on chromosome 13

- RB gene exists in "active" and "inactive" forms
- If active → will stop the advancing from G1 to S phase in cell cycle
- If cell is stimulated by growth factors → inactivation of RB gene → brake is released → cells start cell cycle ...G1 → S→M ...then RB gene is activated again



- Retinoblastoma is an uncommon childhood tumor
- Retinoblastoma is either sporadic (60%) or familial (40%)
- Two mutations required to produce retinoblastoma
- Both normal copies of the gene should be lost to produce retinoblastoma

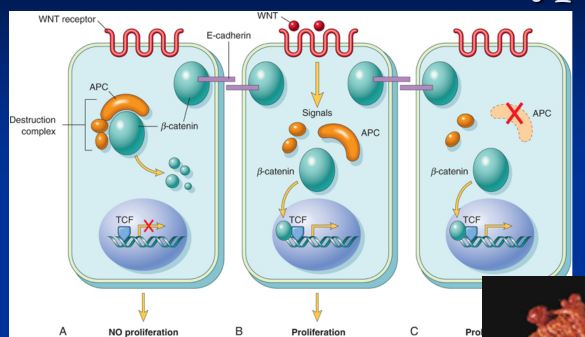
### PATHOGENESIS OF RETINOBLASTOMA Mutation SPORADIC FORM Mutation Somatic cells Germ cells Zygote Somatic cells of child Retinal cells Retinoblastoma of parents FAMILIAL FORM Mutation Mutant Normal RB gene gene

- Transforming Growth Factor- β pathway:
  - TGF- $\beta$  is an inhibitor of proliferation
  - It regulate RB pathway
  - Inactivation of TGF-\( \beta \) lead to cell proliferation

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Mutations in TGF-β pathway are present in :
100% of pancreatic cancers
83% of colon cancers
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- Adenomatous Polyposis Coli β Catenin pathway:
  - APC is tumor supressor gene
  - APC gene loss is very common in colon cancers
  - It has anti-proliferative action through inhibition of β-Catenin which activate cell proliferation
  - Individuals with mutant APC develop thousands of colonic polyps

# Adenomatous Polyposis Coli

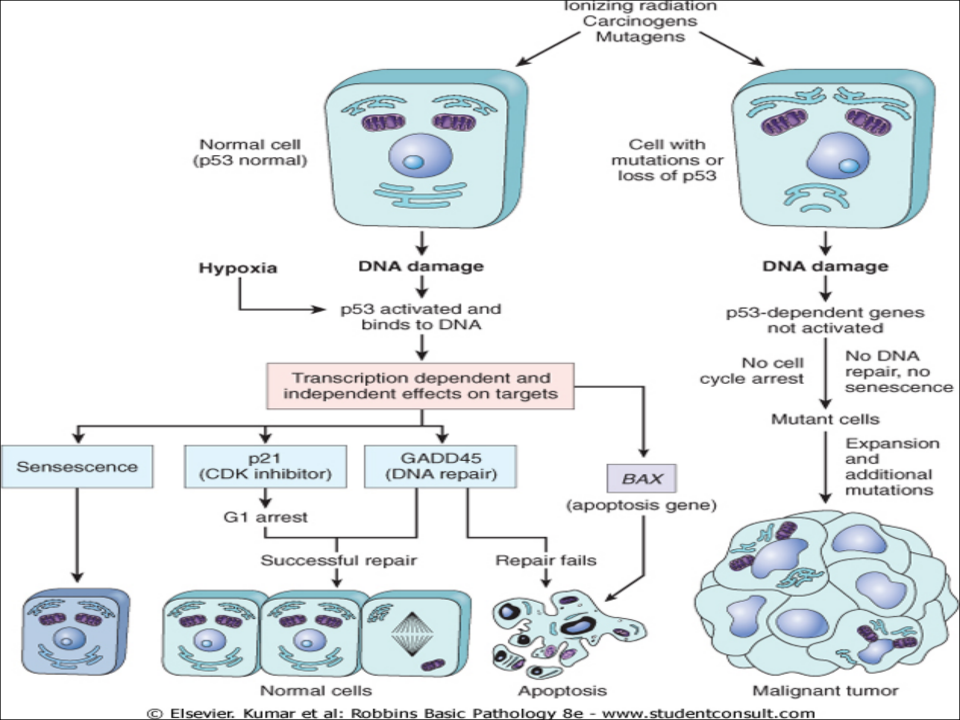


- One or more of the polyps will progress to colonic carcinoma
- APC mutations are seen in 70% to 80% of sporadic colon cancers

- **P**53
  - It has multiple functions
  - Mainly:
    - Tumor suppressor gene (anti-proliferative)
    - Regulates apoptosis

- P53 senses DNA damage
- Causes G1 arrest to give chance for DNA repair
- Induce DNA repair genes
- If a cell with damaged DNA cannot be repaired, it will be directed by P53 to undergo apoptosis

- With loss of P53, DNA damage goes unrepaired
- Mutations will be fixed in the dividing cells, leading to malignant transformation



- P53 is called the "guardian of the genome"
- 70% of human cancers have a defect in P53
- It has been reported with almost all types of cancers: e.g. lung, colon, breast
- In most cases, mutations are acquired, but can be inhereted, e.g : Li-Fraumeni syndrome

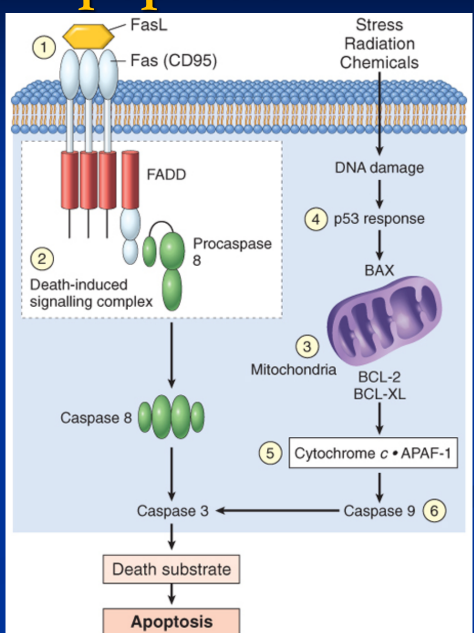
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#### **■** Evasion of apoptosis:

- Mutations in the genes regulating apoptosis are factors in malignant transformation
- Cell survival is controlled by genes that promote and inhibit apoptosis

#### Evasion of apoptosis

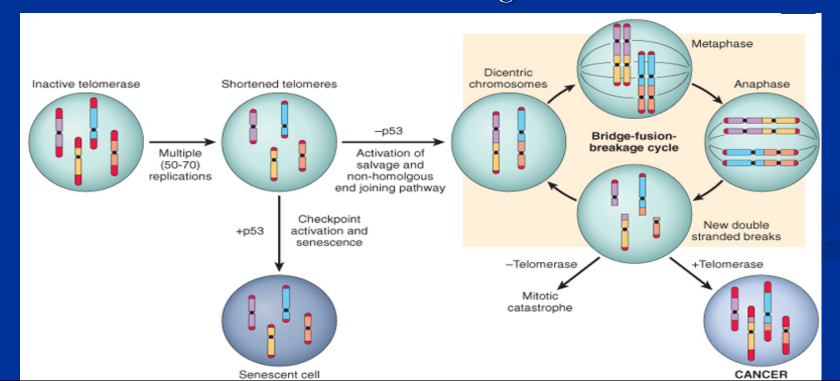
- Reduced CD95 level
  inactivate death —
  induced signaling
  cascade that cleaves
  DNA to cause death →
  tumor cells are less
  susceptible to apoptosis
- DNA damage induced apoptosis (with the action of P53) can be blocked in tumors
- loss of P53 and upregulation of BCL2 prevent apoptosis e.g. follicular lymphoma



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#### ■ Limitless replicative potential:

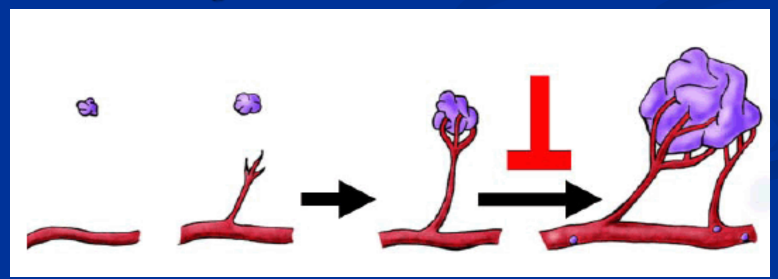
- Normally there is progressive shortening of telomeres at the ends of chromosomes
- Telomerase is active in normal stem cells but absent in somatic cells
- In tumor cells: activation of the enzyme telomerase, which can maintain normal telomere length



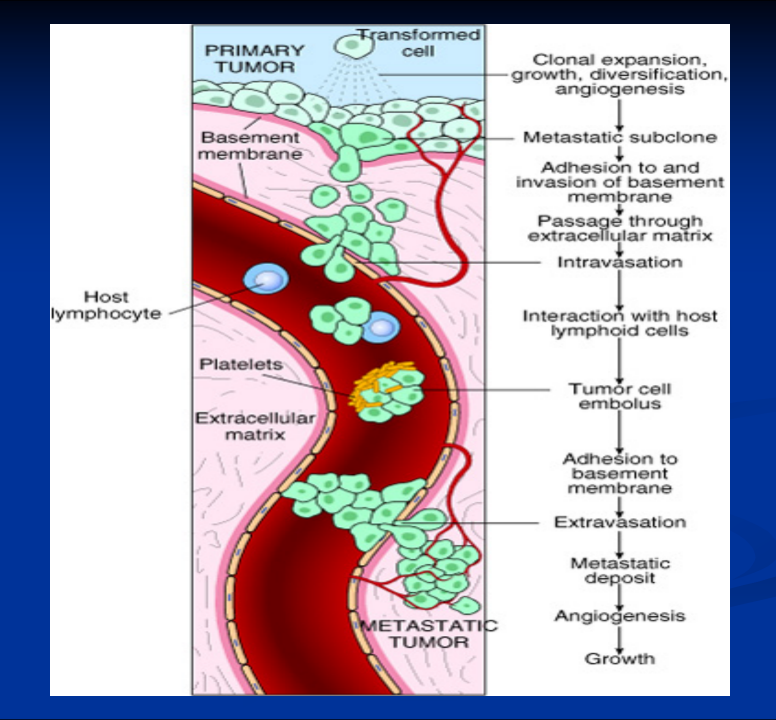
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- Sustained angiogenesis
  - Neovascularization has two main effects:
    - Perfusion supplies oxygen and nutrients
    - Newly formed endothelial cells stimulate the growth of adjacent tumor cells by secreting growth factors, e.g: PDGF, IL-1
  - Angiogenesis is required for metastasis

- How do tumors develop a blood supply?
  - Tumor-associated angiogenic factors
  - These factors may be produced by tumor cells or by inflammatory cells infiltrating the tumor e.g. macrophages
  - Important factors :
    - Vascular endothelial growth factor(VEGF)
    - Fibroblast growth factor



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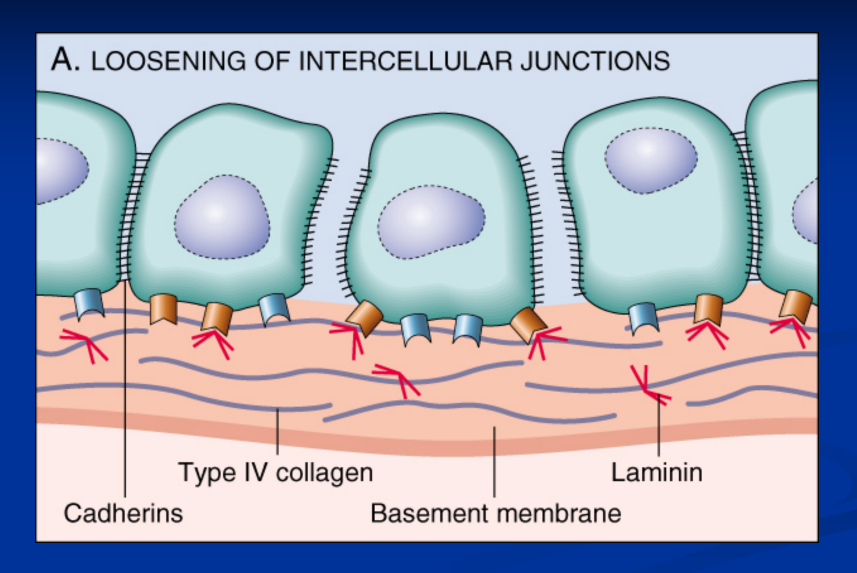


- Ability to invade and metastsize:
  - Two phases:
    - Invasion of extracellular matrix
    - Vascular dissimenation and homing of tumor cells

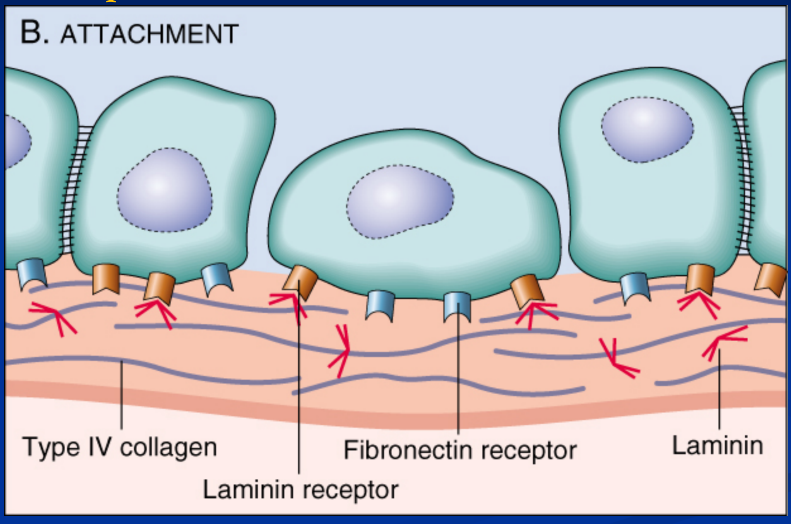
- Invasion of ECM:
  - Malignant cells first breach the underlying basement membrane
  - Traverse the interstitial tissue
  - Penetrate the vascular basement membrane
  - Gain access to the circulation

Invasion of the ECM has four steps:

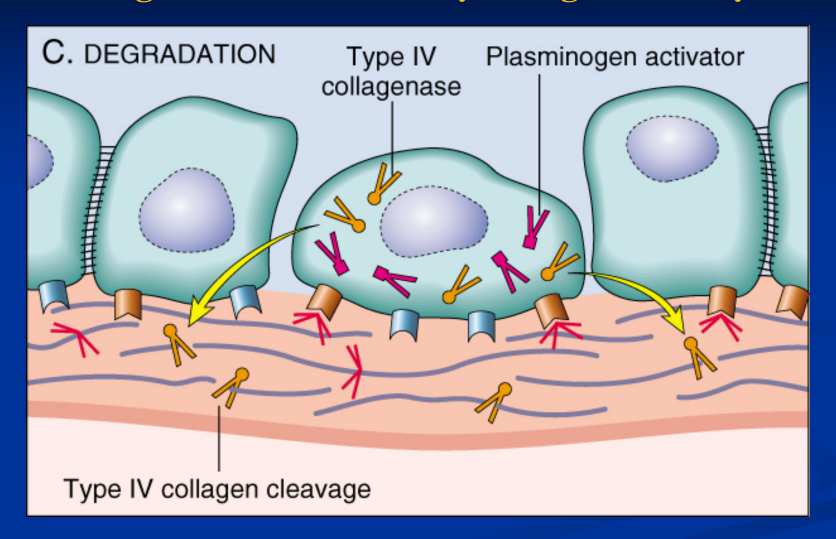
#### 1. Detachment of tumor cells from each other



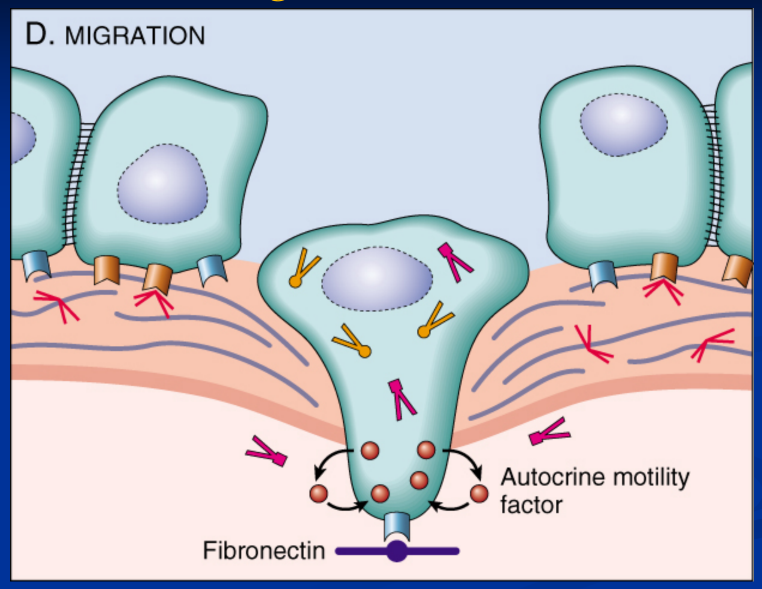
# 2. Attachments of tumor cells to matrix components



#### 3. Degradation of ECM by collagenase enzyme



#### 4. Migration of tumor cells



- Vascular dissemination and homing of tumor cells:
  - May form emboli
  - Most travel as single cells
  - Adhesion to vascular endothelium
  - extravasation

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#### Genomic Instability

- Enabler of malignancy
- Due to defect in DNA repair genes
- Examples:
  - Hereditary Nonpolyposis colon carcinoma(HNPCC)
  - Xeroderma pigmentosum
  - Familial breast cancer

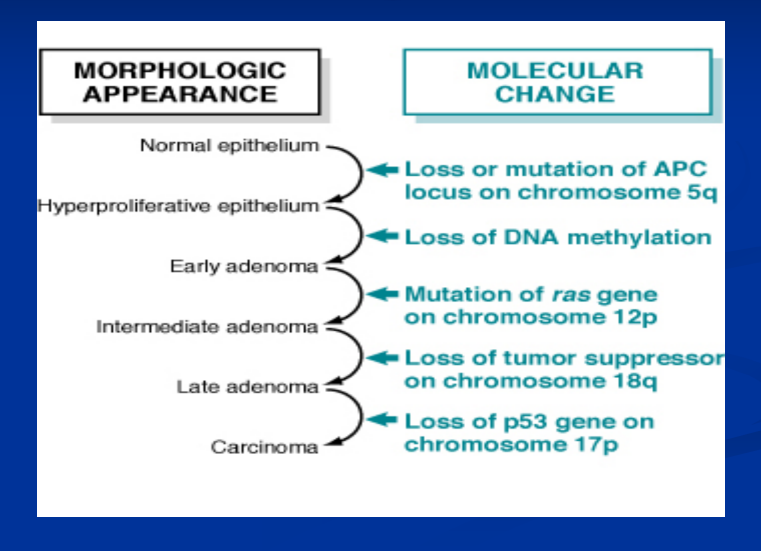
#### Genomic Instability

- Familial breast cancer:
  - Due to mutations in BRCA1 and BRCA2 genes
  - These genes regulate DNA repair
  - Account for 80% of familial breast cancer
  - They are also involved in other malignancies

#### Molecular Basis of multistep Carcinogenesis

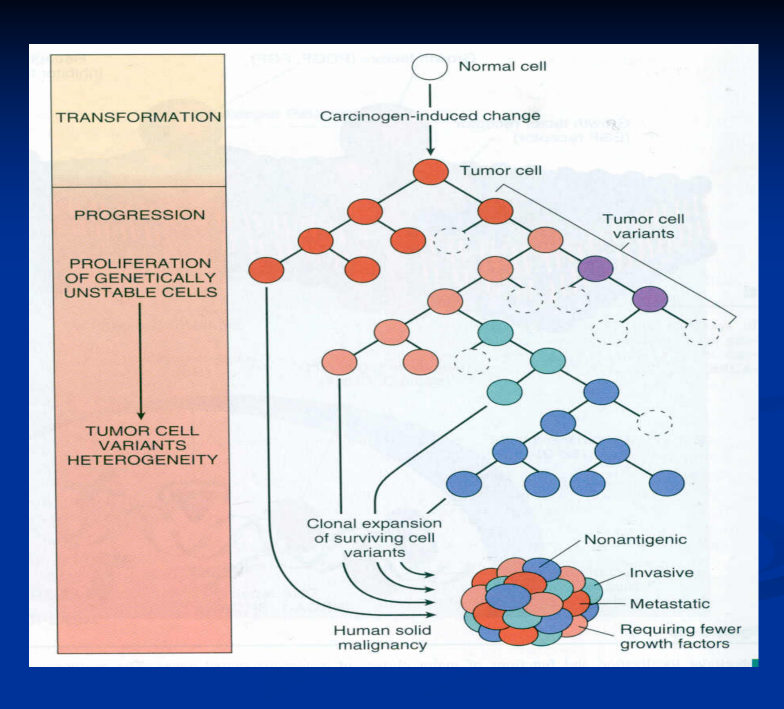
- Cancer results from accumulation of multiple mutations
- All cancers have multiple genetic alterations, involving activation of several oncogenes and loss of two or more tumor suppressor genes

#### Molecular Basis of multistep Carcinogenesis



#### Tumor progression

- Many tumors become more aggressive and acquire greater malignant potential...this is called "tumor progression" ...not increase in size!!
- By the time, the tumor become clinically evident, their constituent cells are extremely heterogeneous



#### Karyotypic Changes in Tumors

- Translocations:
  - In CML: t(9,22) ..." Philadelphia chromosome"
  - In Burkitt Lymphoma : t(8,14)
  - In Follicular Lymphoma: t(14,18)
- Deletions
- Gene amplification:
  - Breast cancer: HER-2

